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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/019,121	05/16/2003	Karin Klokkers	4271-34PUS	8059
7590 04/13/2010 Vincent M. Fazzari Cohen Pontani Lieberman & Pavane			EXAMINER	
			GHALI, ISIS A D	
551 Fifth Avenue Suite 1210 New York, NY 10176			ART UNIT	PAPER NUMBER
			1611	
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

	Application No.	Applicant(s)				
	10/019,121	KLOKKERS ET AL.				
Office Action Summary	Examiner	Art Unit				
	Isis A. Ghali	1611				
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply						
A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS, WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION.  - Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication.  - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication.  - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.7040.						
Status						
1) Responsive to communication(s) filed on 23 December 2009.						
2a) This action is <b>FINAL</b> . 2b) This action is non-final.						
3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.						
Disposition of Claims						
4) Claim(s) 1.3.5-7.10-19 and 21 is/are pending in the application.  4a) Of the above claim(s) is/are withdrawn from consideration.  5) Claim(s) is/are allowed.  6) Claim(s) 1.3.5-7.10-19 and 21 is/are rejected.  7) Claim(s) is/are objected to.  8) Claim(s) are subject to restriction and/or election requirement.						
Application Papers						
9) The specification is objected to by the Examine 10) The drawing(s) filed on is/are: a) acce Applicant may not request that any objection to the c Replacement drawing sheet(s) including the correct 11) The oath or declaration is objected to by the Ex	epted or b)  objected to by the E drawing(s) be held in abeyance. See ion is required if the drawing(s) is obj	e 37 CFR 1.85(a). ected to. See 37 CFR 1.121(d).				
Priority under 35 U.S.C. § 119						
12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).  a) All b) Some * c) None of:  1. Certified copies of the priority documents have been received.  2. Certified copies of the priority documents have been received in Application No.  3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).  * See the attached detailed Office action for a list of the certified copies not received.						
Attachment(s)  1) Notice of References Cited (PTO-892)	4) Interview Summary					
Notice of Draftsperson's Patent Drawing Review (PTO-948)     Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08)     Paper No(s)/Mail Date	Paper No(s)/Mail Da 5) Notice of Informal P 6) Other:	atent Application (PTO-152)				

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#### **DETAILED ACTION**

The receipt is acknowledged of applicants' response filed 12/23/2009 to the non-final office action mailed 07/23/2009.

Claims 1, 3, 5-7, 10-19, 21 are pending and are included in the prosecution.

#### Claim Rejections - 35 USC § 103

- 1. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
  - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- The factual inquiries set forth in *Graham* v. *John Deere Co.*, 383 U.S. 1, 148
   USPQ 459 (1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103(a) are summarized as follows:
  - 1. Determining the scope and contents of the prior art.
  - 2. Ascertaining the differences between the prior art and the claims at issue.
  - Resolving the level of ordinary skill in the pertinent art.
  - Considering objective evidence present in the application indicating obviousness or nonobviousness.
- 3. This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of

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the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

4. Claims 1, 3, 5-7, 10-19, 21 are rejected under 35 U.S.C. 103(a) as being unpatentable over the combination of US 6,303,141 ('141), EP 349 430 ('430), Yanagisawa et al. "Angiogenesis-converting enzyme inhibitors" and Bracht "Transdermal therapeutic system: a review".

### **Applicant Claims**

Claim 1 is directed to matrix-controlled transdermal therapeutic system comprising an active-ingredient-impermeable cover layer, self-adhesive matrix layer, or a plurality of matrix layers of which at least the matrix layer exposed while applying the system is self-adhesive, or one or more matrix layers whose surface remote from the cover layer and intended for adhesion at the application site is coated with an adhesive, the matrix layer(s) comprising at least one ACE inhibitor selected from the group consisting of imidapril, fosinopril, moexipril, perindopril, ramipril, spirapril, cilazapril, benazepril and trandolapril, wherein the inhibitor is in the form of a dicarboxylic acid which is derivatised to form a diester, and removable protective layer.

## Determination of the Scope and Content of the Prior Art (MPEP §2141.01)

US '141 teaches transdermal drug delivery device comprising backing layer, matrix containing 10% ACE inhibitor and Eutanol G as permeation enhancer, and protective release liner. The ACE inhibitor is at least one of benzapril, ramipril or trandolapril which are lipophilic prodrugs of the actual active form of the dicarboxylic acid. Esterification of carboxyl group of ACE inhibitor results in more lipophilic substance (abstract; col.1, lines 16-25; col.2, lines 25-30, 37-43; the claims).

### Ascertainment of the Difference Between Scope the Prior Art and the Claims (MPEP §2141.012)

Although US '141 teaches active forms of ACE inhibitors as dicarboxylic esterified form of the drug, however, the reference does not specifically teach dicarboxylic acid which is derivatised to form **di**ester as claimed by claim 1. US '141 does not teach the cover over the backing layer that is larger than the backing as claimed by claims 14-17.

The cover sheet and its size do not impart patentability to the claims, absent evidence to the contrary.

EP '430 teaches a transdermal system that has improved flux through the skin achieved by using specific salt forms of the drug (page 2, lines 45-50). The transdermal system has a top layer, a layer containing ACE inhibitor including benzaprilat and

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libenzapril, an adhesive layer and protective layer (page 3, lines 40-50). The reference also disclosed on page 3 lines 4-10 the salt forms of the drugs including methane sulphonate and dicarboxylate such as maleate. Claim 3 of the reference teaches lower alkyl dicarboxylate.

Yanagisawa teaches diacid of ACE inhibitor derived from more polar diester was 100 times more patent than one derived from less polar diester (page 423, right column, last paragraph).

Bracht teaches that the use of more lipophilic prodrugs is a strategy to improve the transdermal absorption of molecules. Lipophilic esterification of carboxylic groups can increase the dermal absorption of a drug (page 94, right column, last paragraph).

# Finding of Prima Facie Obviousness Rational and Motivation (MPEP §2142-2143)

Therefore, it would have been obvious to one having ordinary skill in the art at the time of the invention to provide transdermal system for delivery of salts of ACE inhibitors as disclosed by US '141, and replace the salt of ACE inhibitor by lower alkyl dicarboxylate derivatives of ACE inhibitor taught by EP '430. One would have been motivated to do so because EP '430 teaches that transdermal system that having dicarboxylate derivative of ACE inhibitors showed improved flux through the skin. One would reasonably expect formulating transdermal system for delivery of alkyl dicarboxylate of ACE inhibitors at improved flux rates.

Additionally, one having ordinary skill in the art would have been motivated to use more polar diester to derivatise ACE inhibitor dicarboxylate as taught by Yanagisawa. One would have been motivated to do so because Yanagisawa teaches that more polar diester derivatives of diacid form of ACE inhibitor are 100 times potent than those derived from less polar diester. One would reasonably expected formulating transdermal system for delivery of diester derivative of dicarboxylic acid form of ACE inhibitors that is highly potent to treat hypertension effectively.

Furthermore, one having ordinary skill in the art would have been motivated to use the more lipophilic esters of ACE inhibitors in the transdermal delivery as taught by Bracht because Bracht teaches the use of more lipophilic prodrug obtained by lipophilic esterification of carboxylic groups of drugs is a strategy to improve the transdermal absorption of molecules and can increase the dermal absorption of a drug. One would reasonably expect formulating transdermal system for delivery of diester derivative of dicarboxylic acid form of ACE inhibitors that has improved transdermal absorption.

Absent any evidence to the contrary, and based upon the teachings of the prior art, there would have been a reasonable expectation of success in practicing the instantly claimed invention. Therefore, the invention as a whole would have been prima facie obvious to one of ordinary skill in the art at the time the invention was made.

### Response to Arguments

5. Applicant's arguments filed 12/23/2009 have been fully considered but they are not persuasive, and moot in view of the new ground of rejection.

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The examiner hereby responding to applicants' argument against US '141 and EP '430 that remain standing in the current new ground of rejection.

Applicants argue that the Examiner fails to present a proper rationale as to why the claimed invention "as a whole" would have been obvious to a skilled artisan as of the filing date. None of US '141 and EP '430 teaches or suggests a dicarboxylic acid diester derivative of an ACE inhibitor. US '141 is silent about diesters of ACE inhibitors. EP '430 does not teach or suggest any diesters of ACE inhibitors, and instead, discloses ACE inhibitors in their active dicarboxylic acid form or in a salt form. The references in combination fail to provide any basis for the claimed diester derivatives of ACE inhibitors.

In response to this argument, it is argued that one cannot attack the references individually when the rejection is based on combination of the references. US '141 teaches transdermal drug delivery device comprising active form of ACE inhibitor, specifically ramipril or trandolapril. US '141 recognized that esterification of carboxyl group of ACE inhibitor results in more lipophilic substance ACE inhibitors. US '141 however does not teach all the active forms of the ACE inhibitors. EP '430 teaches a transdermal system comprising salt forms of the ACE inhibitors including methane sulphonate and dicarboxylate such as maleate for improved flux through the skin. Therefore, dicarboxylic acid salts of ACE inhibitors were known as taught at the time of the invention. Newly cited reference Yanagisawa teaches diester of diacids of ACE inhibitors as being more potent than other derivatives.

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In any event, applicants' arguments are moot in view of the new ground of rejection. The present invention as a whole is taught by the combined teachings of the prior art.

6. Any inquiry concerning this communication or earlier communications from the examiner should be directed to whose telephone number is (571) 272-0595. The examiner can normally be reached on Monday-Thursday, 6:30 AM to 5:00 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Sharmila Landau can be reached on (571) 272-0614. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

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